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- (54) Glaze compositions.
- (5) A glaze composition comprises 45 to 75% of SiO₂, 0.1 to below 5% of Bi₂O₃, 0.1 to 20% of Al₂O₃, 2 to 20% of B₂O₃, at least one of CaO, MgO, SrO, BaO or ZnO in an amount of 2 to 22%, at least one of Li₂O, Na₂O, K₂O in an amount of 1 to 10%, at least one of La₂O₃, MoO₃ or WO₃ in an amount of 0.1 to 10% and certain other optional ingredients, all percentages being percentages by weight based on the total weight of the glaze composition, the glaze composition having a firing temperature of 950 to 1250°C, and having a coefficient of thermal expansion of 50 to 100 X 10⁻⁷/°C.

The present invention relates glaze compositions and, in particular, to glaze compositions which are substantially free of lead and cadmium and other toxic heavy metal.

For s veral r asons the d velopment of I ad free glazes with the appropriate prop rties t match the propriate so of the ceramic ware to which they are to be applied is a very desirable comm rcial goal. First, occasional episodes of lead poisoning have resulted from the use of improperly formulated and fired lead containing glazes on ceramic ware. Whilst lead containing glazes can be prepared which are safe, and meet current requirements for permissible lead release to food with which they come into contact, the problem of lead poisoning is avoided if lead is avoided. In addition, various pollution controls regarding the use of lead and limits on the content of lead in waste water can be avoided by the use of lead-free glazes.

In the ceramic art a glaze is typically defined as a transparent or opaque glassy coating fired onto a ceramic article or a mixture of ingredients from which the coating is made. Glazes are of two main types, "raw" or "fritted".

"Raw" glazes are typically composed of a mixture of finely ground insoluble beneficiated natural materials - minerals and rocks such as china clay and nepheline syenite. Raw glazes are typically used at high firing temperatures (>1150°C) on substrates such as porcelain (1300°C).

"Fritted" glazes are those where all or part of the ingredients have been prefused and quenched to form one or more frits. The frits are ground and mixed with other constituents (natural materials such as china clay) to formulate the final glaze composition.

Fritting is usually carried out, amongst other reasons, in order to improve homogeneity and to render water soluble or toxic constituents insoluble. Fritted glazes are currently usually used for ceramic ware fired below 1150°C.

The compositions herein described apply to the final composition of the glaze coating, the constituents from which they are formed being partly a matter of availability, economics and choice. It is envisaged that the glaze formulations of the present invention will be utilised as fritted glazes but this is not necessarily always obligatory.

The composition of the glaze is chosen to ensure certain well defined properties such as adhesion to the substrate, a thermal expansion which matches that of the substrate, transparency or opacity, surface finish and texture, and resistance to chemical attack.

Ceramic ware falls into a variety of different categories (Table 1), each with its own characteristic mix of thermal and mechanical properties.

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70.80 \$5.78

980-1080 950-1120

All types, containing frit All types, containing frit

950-1120 single-fire 1100-1150

Berthenuere

Brick

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J	

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			14 17 17 18 18 18 18 18 18 18 18 18 18 18 18 18	,
A Do	Hacult fire (°C)	91010	Clere (iring temperature (°C)	Typical glaze thermal apparation coefficients at 0.70c.1
Continental -	1000	Cleer, leed free, no frit	1350-1400	0\$
Bone Chine	1235	Clear, madium lead or seastless lead free, bigh frit contest.	1020-1100	85-100
Virtect Porei	1250	Gleer, low lead, high frit	1050	60·70
Vitreous senitary ware	Single fire	Opeque, lead free, little or no frit. Often coloured.	1250	60.70
810762876	1000 8 ingle-fire	Various, lead free. No frit	1230	\$
Semi-virrecus mare 1230	1250	Clear, low lead or lead free.	1050	70
Wall cite	1100	Verious, bigh test to test free. Migh frit content. Often textured.	0 9 0 1 - 0 8 4	00.07
ficor tile	1180	some cleaced, therefore nonstip abrasion resistant sizes or engobe needed.	1000-1100	96.70

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Consequently, before one considers any additional requirements as appearance or durability thermomechanical characteristics (glazematuration temperature and the rmal expansion) alone dictates that one glazemiller will not suffice for all substrates. Each type of substrate requires its own type of glaze.

Furthermor , glaz formulations are fr quently tailored not only to match a giv n type of ceramic ware but also the precise conditions under which it will be fired. The glaze will often have been formulated and modified to yield ware of the desired appearance and appropriate physical properties when fired over a specific schedule in a known kiln. It is not uncommon in the ceramics industry to find glazes which are not only unique to a specific manufacturer - but also unique to one of his production kilns. Hence, the ceramics industry uses and has need of a very wide range of glaze formulations.

A durable lead free low melting, low viscosity high refractive index glaze adaptable to a wide range of firing conditions, so characteristic of lead based glazes, has long been sought by the ceramics industry.

The problem has been in finding high refractive index, low melting glazes (≦ 1150°C), without the use of toxic heavy metals such as Pb, Cd or Ba. While elements such as Zr, Ti, La etc can be added to increase the refractive index the accompanying increase in melting temperature or tendency for devitrification has been too great.

In copending European Patent Application No. 91303072.2 we describe a low melting high refractive index(gloss glaze for bone china based on bismuth which solves this problem. Glazes of that invention work well for bone and fine china. The restricted supply and relatively high coat of bismuth however, precludes the use of high bismuth content glazes on a wide spread basis. Such glazes are generally only cost effective on high quality items such as bone china - where top quality is a prerequisite.

A low cost alternative with a wider applicability is required for general useage on white ware. We have found that low bismuth content glazes augumented with a blend of one or more of the elements La, W or Mo can be made to meet this requirement. The glazes of the present invention can be formulated to provide high gloss glazes with thermal expansions and firing temperatures compatible with a wide range of ceramic substrates including; bone or fine china, vitreous or hotel ware and earthenware.

Various lead-free glazes have been described in the art and specific mention may be made of the following prior art.

U.S. Patent No 4285731 discloses frit compositions which are substantially free of lead and cadmium, the frits consisted essentially, in weight percent on the oxide basis, of:

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·: 35 to 47
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                                                           : 5.5 to 9
        B_2O_3
        BaO
                                                           : 24 to 42
        TiO<sub>2</sub>
                                                           : 1.5 to 4
                                                           : 6 to 10
        ZrO<sub>2</sub>
        Li<sub>2</sub>O
                                                           : 1 to 5
        SrO
                                                           : 0 to 8
                                                           : 0 to 5
        MaO
        CaO
                                                           : 0 to 5
                                                           : 1 to 10
        ZnO
                                                           : 0 to 8
        Bi<sub>2</sub>O<sub>3</sub>
        SrO + MqO + CaO + ZnO + Bi<sub>2</sub>O<sub>3</sub> : 0 to 10
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The above frits exhibited a coefficient of thermal expansion between about 65 to 75 X 10-7/°C, a viscosity suitable for firing at about 700° to 950°C, and excellent resistance to attack by acids and bases.

U.S. Patent No 4282035 also describes lead and cadmium-free frits, those frits exhibiting coefficients of thermal expansion between about 52 to 65 x 10⁻⁷/°C, maturing temperatures of about 850° to 1100°C, excellent resistance to attack by acids and alkalis, and consisting essentially, in weight on the oxide basis, of:

```
SiO<sub>2</sub>
                                         : 51 to 60
                                         : 4.5 to 8
        B<sub>2</sub>O<sub>3</sub>
        BaO
                                         : 0 to 13
        SrO
                                         : 0 to 18
                                         : 6 to 30
        BaO + SrO
                                         : 4 to 8
        ZrO_2
        Al_2O_3
                                         : 5 to 8
        Li<sub>2</sub>O
                                         : 0 to 4
                                         : 0 to 5
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        Na<sub>2</sub>O
                                         :0t 5
        K<sub>2</sub>O
        Li_2O + Na_2O + K_2O
                                         : 1 to 5
                                         :0t 6
        MgO
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CaO : 0 t 12 Bi₂O₃ : 0 to 10 MgO + CaO + Bi₂O₃ : 0 to 20

U.S. Patent No 4554258 describes lead, cadmium and arsenic fr e glass frit compositions which consist ssentially of Bi₂O₃, B₂O₃, SiO₂ with R₂O being 2 to 8% and RO b ing 0 to 9% by weight.

U.S. Patent No 4590171 also describes lead and cadium free frits consisting essentially, in weight percent on the oxide basis of

Li₂O : 3 to 4 Na₂O : 0.75 to 3 BaO : 3.5 to 9.5 B_2O_3 : 14 to 17.5 : 6.75 to 8.75 Al_2O_3 SiO₂ : 48 to 55 ZrO_2 : 6.75 to 10.5 F : 3 to 4

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Finally, U.S. Patent No 4892847 discloses lead free glass frit compositions consisting essentially of SiO₂, Bi₂O₃, B₂O₃, alkali metal oxide and ZrO₂/TiO₂ in appropriate concentrations.

There are four essential criteria which must be demonstrated by glazes, and a fifth characteristic which is essential when a high gloss glaze is required.

First, the firing or glazing temperature of the glaze must not exceed the temperature at which the ceramic body being coated thermally deforms.

Second, the coefficient of thermal expansion of the glaze must be compatible with that of the ceramic body being coated to preclude crazing and/or spalling; the glaze preferably having coefficients of thermal expansion somewhat lower than that of the substrate such that, when the ceramic body is cooled the fired coating will form a surface compression layer.

Third, the glaze must possess excellent resistance to attack by acids and bases since corrosion of the coating can result in the loss of gloss, the development of haze and/or iridescence, the formation of porosity, or other defects deleterious to the appearance or physical character of the coating.

Fourth, in the case of transparent glazes, the glaze must maintain good glass stability and must not devitrify to any substantial extent during the firing.

Fifth, where a glossy appearance is desired, the refractive index of the glaze must be high.

We have now developed lead-free glaze compositions which can be applied to a wide range of ceramic substrates including bone or fine china, vitreous ware and earthenware.

Accordingly, the present invention provides a glaze composition which is essentially free from lead and cadmium and which comprises the following components:

	sio ₂	:	45	to	75%	
	Bi ₂ 0 ₃	:	0.1	to	below	5%
40	Al ₂ 0 ₃	:	0.1	to	20%	
	B ₂ O ₃	:	2	to	20%	
	at least one of					
45	Ca0, Mg0, Sr0, Ba0 or Zn0	:	2	to	228	
	at least one of					
	Li ₂ 0, Na ₂ 0, K ₂ 0	:	1	to	10%	
	at least one of					
50	La_20_3 , $Mo0_3$, $W0_3$:	0.1	to	10%	
	TiO ₂	:	0	to	10%	
	Zr02	:	0	to	10%	
55	P ₂ 0 ₅	:	0	to	5%	
	V ₂ 0 ₅	:	0	to	18	
	fluoride ions	:	0	to	5≹	

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all p rc ntag s b ing p rc ntag s by w ight bas d n the total w ight of th glaz composition, the glaz composition containing La_2O_3 , MoO_3 and/or WO_3 in a total amount of less than 10% by weight, with the proviso that th amount of ach of WO_3 and MoO_3 is less than 5% by weight, and with th proviso that BaO is not contained is an amount of more than 2% by weight, the glaze composition having a firing temperature of 950 to 1250°C, and having a coefficient of thermal expansion of 50 to 100 X 10-7/°C.

The glaze compositions of the present invention preferably comprise from 1.0 to 4.5% by weight of Bi_2O_3 , more preferably 2.0 to 4.0% by weight of Bi_2O_3 , from 50 to 65% by weight of SiO_2 , from 7 to 15% by weight of AI_2O_3 , more preferably from 8 to 14% by weight of AI_2O_3 , and from 5 to 14% by weight of Bi_2O_3 , more preferably from 7 to 12% by weight of Bi_2O_3 .

The glaze compositions of the present invention contain at least one of La_2O_3 , MoO_3 or WO_3 in the amounts as specified above. When the additive is MoO_3 it is preferably used in an amount of less than 2% by weight, preferably less than 1% by weight as amounts above 2% by weight will tend to render the glazes opaque. When the additive is WO_3 it is preferably used in an amount of less than 2% by weight, preferably less than 1% by weight as amounts above 2% by weight will tend to render the glazes opaque. When the additive is La_2O_3 it is preferably used in an amount of less than 5% by weight. The total amount of La_2O_3 , MoO_3 and WO_3 is preferably 0.1 to 8.0% and more preferably in the range of from 0.5 to 5.0% by weight.

Due primarily to the use of MoO₃ and WO₃ many of the glaze formulations of the invention have an increased tendency towards liquid Immiscibility which can result in opalescence or opacity.

Hence, when a transparent glaze is required appropriate and standard steps within the scope of the present invention can be taken to minimise this. It is, for example, known in the art that increasing amounts of MoO_3 and WO_3 promote immiscibility, as can P_2O_5 , whereas Al_2O_3 suppresses it.

In general, the trend is that the higher the lonic field Strength (Z/r) of an ion the more likely it is to promote immiscibility. Hence, such elements ought to be minimised to an optimimum level. Lithium, for example, is more likely to induce liquid immiscibility than are Na or K and so should not be used in excess.

It has also been found that some formulations are sensitive to the precise firing conditions. In general, higher temperatures and faster cooling inhibit liquid immiscibility and result in clear glazes. This in fact makes such glazes well suited to the fast fire schedules now being introduced into the tableware industry.

As is general practice, the precise formulations of the glazes of the present invention should be precisely matched not only to the intended substrates but also to the kiln firing cycles (and atmospheres) to be used in order to obtain the desired appearance.

In order to provide a high gloss the glaze of the present invention should preferably have a refractive index of at least 1.45.

The glass frits which may be used to form the glazes of the present invention can be prepared by mixing together the oxide producing materials, charging the mixture of materials into a glass melting furnace at a temperature sufficiently high to produce the fused glass and then fritting the glass by pouring it into water, or by passing it through water cooled rolls. It may be preferred to carry out the fritting operation in an oxidising atmosphere, or to include an oxygen rich component in the mixture which is melted and fused. The frit may be ground into a powder by conventional grinding techniques.

Though methods exist for applying glazes in dry or even molten form, the conventional method of application for bone china is in the form of a fine ground water based slurry or slip with the item being coated by either dipping or spraying. This glaze slip might consist, in addition to the carrier medium-water, exclusively of a single ground frit, or a mixture of many materials including frits, minerals, insoluble manufactured chemicals as well as minor quantities of rheological modifying agents. These latter constituents may include such things as floculants, deflocculants, binders, wetting agents, antifoaming agents and for identification purposes even organic dyes. The glaze may optionally contain one or more pigments.

The mode of application of the glaze to the ceramic article is not seen as central to this invention, any feasible method is permissible.

The precise blend and form of the constituents from which the final glaze is formed may be changed without substantially departing from the invention intended to be defined herein. The description being merely illustrative of embodiments of the principle of the invention and not limiting it.

The glazes of the present invention can be formulated to have a glazing or glost temperature in the range of from 950 to 1250°C in order to make them suitable for use with bone or fine china vitreous ware and earthenware. By the term "glazing temperature" as used herein is meant a temperature at which the glaze melts and flows sufficiently to produce a smooth uniform homogen ous coating on the ceramic body being coated the rewith. The glazing temperature is also sometimes referred to as the firing or glost temperature.

The glazes of the pr s nt inv ntion compris at I ast one of CaO, MgO, SrO, BaO or ZnO in an amount of from 2 to 22% by weight, in total. Mixtur s of these oxides may advantag ously b used. Barium is pr ferably absent but if us d the amount of BaO in the glazes of th pr sent invention is k pt below 2% by weight in view

of its t xicity. The glazes of the present invention alse comprise at least one of Li₂O, Na₂O or K₂O in an amount from 1 th 10% by weight, in total. The amount of Li₂O is preferably 1 so than 4% by weight. When ZnO is used in the glazes of the present invention it is preferably used in an amount of below 2% by weight.

The glazes of th present invention may also comprise the various optional ingr dients listed above. Additions of ZrO₂ and/or TiO₂ are b neficial since ZrO₂ will improve the resistance of the glaze to alkali and detergent solution attack, whilst the addition of TiO₂ will improve the resistance of the glass to acid attack. The total addition of ZrO₂ and/or TiO₂ is preferably less than 4% by weight.

The present invention also includes within its scope a method of glazing a ceramic body, which method comprises coating a ceramic body with a glaze composition as hereinbefore described and firing the coated ceramic body at a temperature of 950 to 1250°C.

Preferably the ceramic body coated according to the method of the invention is vitreous ware or earthenware

The present invention will be further described with reference to the following examples.

15 Examples 1 and 2

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The glazes of Examples 1 and 2 were prepared by the following general procedure.

A glass of the appropriate composition was melted at approximately 1300°C, frit quenched, ground to a powder and the mixed with 12% china clay by ball milling in water to produce a glaze slip which, on firing, yielded a glaze composition as given in Table 2.

Both trials followed the same procedure. After coating the test piece (bone china plate) with the glaze slip and drying, they were fired by heating at 5.9°C/min to 1100°C, held for 45 minutes and cooled at 5.9°C/min to room temperature. A glossy glaze was obtained.

25		Table 2	
		1	2
	sio_2	59.61	56.92
30	Al ₂ 0 ₃	9.92	9.47
	Ca0	12.37	6.42
	Sr0	0.99	12.10
	к20	2.25	3.26
35	Na ₂ 0	2.97	0.51
	Li ₂ 0	-	0.12
	B ₂ O ₃	4.94	5.82
40	La ₂ 0 ₃	2.98	1.00
	Bi_20_3	2.98	1.98
	Mo03	_	0.40
	wo ₃	0.99	1.00
45	zro ₂	-	1.00

Examples 3 to 88

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The glazes of these Examples were prepared by the following general procedure.

The components were mixed in a mortar and pestle and fused at 1350° C for 1^{1} /4 in a platinum crucible. The glass produced was quenched and the frit initially hand ground, followed by ball milling in water for 4 hours in a ceramic pot with an Al_2O_3 milling medium (average particle size of final milled frit was approximately 6 micrometr s). The slurry was dried and then 12% china clay added with a further 15 minute milling period to yield a glaze slip which on firing gav the compositions as d tail d in Tabl 3 b low.

Approximat ly 8g of the glaze was spray d onto the chos in ceramic substrat or substrat is sill cted from 6 inch diameter bon ichina plates, arthenwar plates or hotel ware plates and thin fired as described below.

The arthenware and hot I ware plates w r glost fired employing a ramp rate of 3°C per minute to 1100°C,

a 2 hour holding tim and a cooling rate of 5°C p r minute. A variation was mad for firing onto bon china plat s wh n th holding temperature was 1080°C for 2 h urs.

The glazes were fired onto the following substrat s and in all cases produced a good gloss.

Bon China only

- Examples 3 to 5.

Bone China and earthenware

- Examples 6 to 8.

Earthenware only

- Examples 9 to 23, 34 to 36, 40, 44, 45, 47, 48, 50, 68,

70 to 76.

Hotel ware only

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- Examples 24 to 33, 42, 43, 56, 58, 62, 63, 66, 67, 83

to 88.

Earthenware and Hotel ware

- Examples 37 to 39, 41, 46, 49, 51 to 55, 57, 59 to 61,

64 and 65.

The glazes were also tested by means of 'Flow Trials'. This is a standard test used in the industry to give a measure of the melting and flow (viscosity) characteristics of a glaze the appearance of which also gives a preliminary indication of glaze stability (tendency for crystallization and phase immiscibility).

For the flow trials, 1.5gm of glaze powder was dry pressed into a disc and placed in a circular slot at the mouth of a flow channel in a flow trial slab. The slab was mounted at an angle of 33 degrees and fired at a ramp rate of 3°C per minute to 1100°C, with a 2 hour holding time and a cooling rate of 5°C per minute.

In the Examples given in Table 3 the following abbreviations are given for the appearance of the glaze in the flow trial:

O.W. = opaque white

OP = opal

5 S.O. = slight opalescence

V.S. = very slight opalescence

T = transparent

Some flow trials were also performed at 1200°C using the same ramp rates but a shorter holding time of 1 hour.

At this higher temperature the following abbreviations were used.

- = transparent
- o = very slight opalescence

Some flow trials were also performed at 1130°C with a ramp rate of 9°C per minute to 1130°C, a 45 minute holding time and a cooling rate of 9°C per minute to mimic a 'fast fire' schedule. At this temperature the following abbreviation was used.

 Δ = transparent

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5																						
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15			11	59.25	11.33	13.22	7.13	2.08	2.00	2.00	1.00	1.00	1.00			ı		OP			6.17	
,,			10	59.25	11.33	13.22	7.12	4.08	1.00	1.00	1.00	1.00	1.00	1	1	•	1	8.0.			0.9	
20			o	57.42	10.47	11.10	90.9	5.03	0.50	0.50	2.97	2.97	0.99	,	,	1.00	0.99	90				
25			60	56.75	11.33	11.22	6.12	5.08	0.50	0.50	3.00	3.00	1.00	0.30	ı	1	1.00	0P*			8.34	
30		TABLE 3	7	58.75	11.33	13.72	2.12	4.08	1.00	1.00	0.50	1.25	1.25	ı	1	1	9.00	*40			7.86	
			v	58.0	10.58	11.22	6.12	5.08	0.50	0.30	3.00	3.00	1.00	,	ı	,	1.00	v.s.			8.8	•
35			K O	90.09	6.32	5.58	13.88	3.36	2.32	,	1.68	1.68	0.56	1	•	ı	0.12	8.0.			8.94	
40			4	60.76	6.23	5.58	13.88	3.36	2.32		3.35	3.35	1.12	ı	•	t	0.12	MO			ι	
45			m	57.14	5.66	6.52	7.18	0.56	3.44	0.12	2.22	1.12	1.12	0.44	1.12	1	13.58	*40			9.0	
50			Example No.	5102	A1203	B203	Cao	Na ₂ 0	K20	L120	B1203	La203	WO3	No03	Zro2	Zn0	Sr0	Appearance	in flow trial	at 1100°C	Thermal	Expansion (X 10 ⁻⁶ /°C)

										•													
10																							
15		20	58.25	11.33	13.22	9 0	9 6	8 5	1.00	1.00	2.00	1	•	,	1	•	3			6.97			
15		19	59.25	11.33	13.22	7.12	80.4	J. 00	9 6	} '	2.00	; '	,		•	ı	3			6.57			
20	,	18	60.26	11.33	9.22	7.12	4.08	1.00	1.00	4. ·		00.1	ı	•		ı	. ;			OR S			
25		17	60.26	11.33	9.33	7.13	4.08	1.00	1.00	66.4	7.00	1	1		•	1	•	F			9.		
•	continued	16	56.25	11.33	11.22	7.12	4.08	1.00	1.00	3.00		2.00			ı			K.		,	6.63		
30	TABLE 3 - Continued	15							1.00									оъ			6.13		
35	E4	14	-						1.00									0.W.			7.9		
40		13	κ π		11.22	7.13	4.08	1.00	1.00	3.00	3.00	3.00	ı	•	1	ı		O.W.			7.13		
		12																0.W.			6.87		
45		<u>.</u>	-	., •		•													trial	ပွ		ro co	(20/
50		Example No.		8102	A1203	6203	200	Na20	1.1.0	B1,0,1	[a,0]	W03	Moor	zro,	Zu0	gro	É	Appearance	in flow trial	at 1100°C	Thermal	Expansion	$(x 10^{-6}/^{0C})$

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10																							
		29	63.55	11.33	9.50	7.12	2.75	2.75	0.30	1.00	0.50	0.50	0.50	•	•	•	1	×.0	: !		7.06		
15		28	63.55	11.33	9.50	7.12	2.00	2.00	2.00	1.00	0.50	0.50	0.50	•	ı	1	ı	, M. O			6.43		
20		27	59.00	12.58	9.47	7.12	4.25	2.58	2.00	2.00	1.00	,	•	1	ı	•	•	F	ı		١		
25		52	62.50	11.00	9.00	6.00	3.00	2.00	1.50	1.00	1.00	1	ı	0.50	1	2.00	0.50	E	,		7.00	•	
	to to	25	63.00	11.00	9.00	6.00	3.00	2.00	1.50	1.00	1.00	•		0.50	•	2.00		8.0.			69.9	-	
30	ARES 3 - Continued	34	60.26	11.33	9.22	7.12	1.58	4.00	0.50	4.99	1.00	•	ı	1	,	,	,	8.0.			6.91		
35	F	23	60.26	11.33	9.22	7.12	1.58	4.00	0.50	4.99	1.00	,	ı	•	•	1	1	8.0.			6.80		
40		22	57.25	11.33	13.22	7.12	4.08	1.00	1.00	1.00	1.00	1.00	1	ı	•	•	2.00	90			•		
		21	57.25	11.33	13.23	7.12	4.08	1.00	1.00	1.00	1.00	1.00	•	2.00	•	•	•	V.S.			•		
45		Example No.	S102	203	33		0.	•	0.	t ₀ 3	103	_	5	2		_		earance	flow at	200	rmal	anston	10-6/oc)
50		Bxa	sic	A12	B2C	Ca	Naz	K20	17	812	ž	WO3	Ko	2r0	2n0	3r 0	Tio	App	Į,	110	The	Exp	č

40 45		Example No. 30 31	02 63.55 63.55	11.33	9.6	7.13	2.75	0.50	2.75	1.00	0.50	0.50	0.50	ı	1	ı	•	ŧ	at 1100°C	lermal
35	н	32	5 64.75	-																9.9
30	TABLE 3 - Continued	33	64.45	11.33	9.50	7.12	2.00	2.00	2.00	1.00	0.20	0.20	0.20	1	•		,	s.0.		6.37
	ntinued	34	63.45	11.33	9.50	7.12	4.00	0.50	1.50	1,00	0.20	0.20	0.20	1.00	ı	•	•	ď		٠
25		35	63.75	11.33	9.50	7.12	2.00	2.00	2.00	1.00	0.10	0.10	0.10	1.00	1	•	•	ďO		•
20		36	63.00	11.00	9.00	00.9	4.00	1.00	1.50	1.00	0.20	0.20	0.20	0.50	•	3.00	ı	0 b		•
15		37	58.75	11.33	14.22	7.12	4.08	1.00	1.00	0.50	1.00	ı	1.00		1	•	1	V.8.		6.74
		38	63.00	11.00	00.6	6.00	3.00	2.00	1.50	0.50	0.50	ı	0.50	1.00	•	2.00		E		69.9
10																				
5																				

4 5	45	40	35	30	00	25	20	15	15	10	5
			ĺ								
			3	TABLE 3 - COULTINGE	Desturant						
Example No.	36	40	41	42	43	4	4.5	46	47		
8102	61.50	57.45	63.75	63.00	63.00	60.88	63.75	58.25	56.25		
A1203	11.00	11.33	11.33	11.00	11.00	11.00	11.33	11.33	11.33		
B203	9.00	13.22	9.22	9.00	00.6	9.00	9.33	9.22	11.22		
CaO	8.00	9.00	7.12	6.00	6.00	7.12	7.12	7.12	7.12		
Na20	3.00	2.00	2.08	3.00	3.00	4.00	3.08	6.08	6.08		
K20	2.00	2.00	2.00	2.00	3.00	2.00	2.00	•	1		
L120	1.50	2.00	2.00	1.50	1.50	0.50	2.00	1	•		
B1203	0.50	0.50	1.00	1.00	1.00	3.00	1.00	5.00	3.00		
La203	0.50	1.00	1.00	1.00	1.00	1.00	1.00	3.00	5.00		
WO3	•	ı	•	•	•	•	,	•	1		
No03	0.50	0.50	0.50		•	0.50	0.50	ı	ı		
2012	0.50	0.50	ı	0.50	0.50	ı	•	ı	1		
2n0	1.00	0.50	,	•		1.00	•	,	1		
8r0	1.00	ı		2.00	2.00	ſ	•	ı	•		
T102	•	ı	,		•	1		•	1		
Appearance	H	Ħ	v.s.	V.S.	v.s.	8.0.	9.0.	E	oP		
in flow trial											
at 1100°C											
Thermal	•	ı	•	1	·		1	ı	1		
Expansion					•						
(X 10-6/0C)											

45 50		40	35	30		25	20	15		10
		,	TA	TABLE 3 - Continued	ntinued					
Example No.	8	64	20	51	. 52	53	54	55	56	
5102	. 38.25	56.25	56.25	56.25	59.26	59.25	59.26	55.76	57.76	
A1203	11.33	11.33	11.33	11.33	11.33	11.33	11.33	11.33	11.33	
B ₂ 0 ₃	11.22	9.23	11.22	11.22	8.72	10.72	9.22	11.22	9.22	
Cao	7.12	7.12	7.12	7.12	7.12	7.12	7.13	7.13	7.12	
Na ₂ 0	80.9	6.08	4.08	3.08	3.08	3.08	4.08	4.08	4.08	
K20	ı	•	1.00	2.00	2.00	2.00	1.00	1.00	1.00	
L120	•	•	1.00	1.00	1.50	1.50	1.00	0.50	0.50	
Bi203	3.00	5.00	3.00	3.00	4.99	3.00	4.99	4.99	4.99	
La203	3.00	5.00	2.00	5.00	1.00	1.00	1.00	3.00	3.00	
W03		1	1	•	•	•			t	
Mo03	•	.	ı	1	ı	ı	•		ι	
Zr02	•	•		•	0.50	0.50	0.50	0.50	0.50	
Zno	•	,	1	ı	0.50	0.50	0.50	0.50	0.50	
Sro	•	1	1		ı	•	ı	ı	•	
T102	ı		•	•		•	1		ı	
Appearance	V.S.*, A	o B	ďO	ďO .	Ħ	E	E+	e	H	
in flow trial										
at 1100°C								;		
Thermal	ı	,		•	•	,	7.37	8.03	ı	
Expansion (X 10 ⁻⁶ /°C)										

50	45	40	35		30	25	20	15	15	10
		•, •,	ä	TABLE 3 - Continued	ntinued					
Example No.		28	86	09	61	62	63	. 49	65	
S10 ₂	55.76	57.26	58.76	56.76	55.76	55.76	57.76	61.76	59.70	
A1203	13.33	13.33	13.33	13.33	13.33	13.33	13,33	11.33	11.33	
B203	9.23	9.23	7.22	7.22	9.32	7.22	7.22	7.22	7.22	
CBO	7.12	7.12	7.12	7.12	9.12	9.12	9.12	7.12	9.12	
Nago	4.08	4.08	4.08	4.08	4.08	4.08	4.08	4.08	4.08	
K20	1.00	1.00	2.00	2.00	1.00	1.00	1.00	1.00	1.00	
1,20	0.50	1.00	0.50	0.50	0.50	0.50	0.30	0.50	0.50	
B1203	4.99	4.99	4.99	4.99	4.99	4.99	4.99	4.99	4.99	
La203	3.00	1.00	1.00	3.00	1.00	3.00	1.00	1.00	1.00	
WO3	1	1	1		ı	•	ı	1	٠	
Mo03	ı	•	•		•	•	•	•	ı	
2r02	1	0.50	0.50	0.50	0.50	0.50	0.50	0.50	0.50	
Zno	:	0.50	0.50	0.50	0.50	0.50	0.50	0.50	0.50	
8r0	1	•	•	•				•	•	
Mo03	ı	1	•	•	•		•	•	1	
T102	ı		1	1		1	1	ı	1	
Appearance	Ħ	H	H	E	E	E	£	8.0.	E	
in flow trial						ı	ı		ı	
at 1100°C										
Thermal	7.06	•	7.06	7.77	7.49	7.46	7.31	7.03	7.49	
Expansion								1		
(X 10-6/0C)										

50	4 5	40	35	30		25	20	15	10	5
	:		ĺ							
			T.	IABLE 3 - CONTINUES	ntinued					
Example No.	75	76	7.7	78	62	80	81	82	83	
8102	57.50	57.77	30.00	50.25	60.25	65.37	54.90	68.01	59.01	
A1203	10.64	10.69	20.00	11.33	11.33	11.33	13.00	11.00	11.00	
B203	13.35	13.42	14.00	13.22	5.22	13.22	20.00	2.00	5.00	
Ca0	69.9	6.73	20.00	7.12	7.12	2.00	7.00	7.00	5.00	
Na ₂ 0	5.71	5.74	8.00	80.9	4.08	6.08	4.00	9.00	4.00	
K20	ı		1			,	•		4.00	
L120	ı		ı	ı		•	ı		1.00	
B1203	2.82	2.83	1.00	1.00	1.00	1.00	0.10	4.99	4.99	
La ₂ 0 ₃	2.83	2.83	5.00	1.00	1.00	1.00	1.00	1.00	2.00	
WO ₃	•	ı	1	•	ı	ı	ı	•	ı	
Mo0 ₃	•	ı		1.		•	•	1	•	
Zr02	ı	•	2.00	ı		•		•	1	
2n0	1	1	1		10.00		•	ı	2.00	
Sro	•	ı	•		1	•	ı			
T102	1	•	. •	10.00	•	•	•			
V205	0.47	1	•			•	•	•	•	
Mg0	1	•	•	1		•	•	•	2.00	
Appearance	d O	do	£	H	8.0.	8.0.	E	8.0.	F	
Flow Trial										
at 1100°C										
Thermal	ł		1		1	,			1	
Expansion (X 10 ⁻⁶ /°C)										

5 10 15 20 25 61.10 6.00 6.50 2.00 1.00 6.00 1.00 1.00 3.00 3.00 3.00 30 11.00 7.00 2.00 5.00 3.00 1.00 4.99 1.00 0.50 0.50 1.00 35 TABLE 3 - Continued 1.50 0.50 0.50 1.50 1.00 1.00 40 5.00 112.00 14.00 2.00 1.00 3.00 1.00 2.00 2.00 2.00 45 MOO3
ZEO2
ZEO
SEO
MGO
TIO2
V205
Appearance
Flow Trial
at 1100°C
Thermal
Expansion
(X 10^-6/°C) Example No.

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Some glaze slip was fir d and melted in a ceramic mould using the standard 1100°C firing cycl to form a glass bar.

Fragments of this bar w re coars ground using a p stle and mortar. R fractive index m asur ments were made on the r sultant powder samples with an optical microscope using th standard B ckke Lin Oil immersion technique.

The results are shown in Table 4 below:

TABLE 4

10	Glaze Composition	Refractive Index
	11	1.520 ± 0.005
15	13	1.530 ± 0.005
	15	1.530 ± 0.005
	16	1.530 ± 0.005
20	17	1.530 ± 0.005
	54	1.525 ± 0.005
	55	1.525 ± 0.005

Claims

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 A glaze composition which is essentially free from lead and cadmium and which comprises the following components:

	sio ₂	:	45	to	75%	
	Bi ₂ 0 ₃	:	0.1	to	below	5%
35	Al ₂ 0 ₃	:	0.1	to	20%	
	B ₂ 0 ₃	:	2	to	20%	
	at least one of					
40 .	CaO, MgO, SrO, BaO or ZnO	:	2	to	228	
	at least one of					
	Li ₂ 0, Na ₂ 0, K ₂ 0	:	1	to	10%	
	at least one of					
45	La ₂ 0 ₃ , MoO ₃ , WO ₃	:	0.1	to	10%	
	TiO ₂	:	0	to	10%	
	ZrO ₂	:	0	to	10%	
50	P ₂ 0 ₅	:	0	to	5%	
	v ₂ o ₅	:	0	to	18	
	fluoride ions	:	0	to	5%	

all p rcentag s being percentages by w ight bas d on the total weight of the glaze composition, the glaze composition containing La₂O₃, MoO₃ and/or WO₃ in a total amount of less than 10% by w ight, with the proviso that the amount of each of WO₃ and MoO₃ is less than 5% by w ight, and with the proviso that BaO is not contained is an amount of more than 2% by weight, the glaze composition having a firing tem-

- p ratur of 950 t 1250°C, and having a coefficient of thermal expansion f 50 to 100 X 10-7/°C.
- 2. A glaze composition as claim d in claim 1 which comprises from 1.0 to 4.5% by weight of Bi₂O₃.
- 3. A glaze composition as claim d in claim 1 or claim 2 which comprises from 50 to 65% by weight of SiO₂.
 - 4. A glaze composition as claimed in any one of the preceding claims which comprises from 7 to 15% by weight of Al₂O₃.
- 5. A glaze composition as claimed in any one of the preceding claims which comprises 0.1 to 2% by weight of MoO₃.
 - 6. A glaze composition as claimed in any one of the preceding claims which comprises 0.1 to 2% by weight of WO.
- 7. A glaze composition as claimed in any one of the preceding claims which comprises 0.1 to 5% by weight of La₂O₃.
 - A glaze composition as claimed in any one of the preceding claims which contains a total amount of La₂O₃, MoO₃ and WO₃ in the range of from 0.1 to 8.0% by weight.
- A glaze composition as claimed in any one of the preceding claims which comprises from 5 to 14% by weight of B₂O₃.
 - 10. A glaze composition as claimed in any one of the preceding claims which fires at a temperature range of from 950 to 1250°C.
 - 11. A glaze composition as claimed in any one of the preceding claims which has a refactive index of at least 1.45.
 - 12. A glaze as claimed in any of the preceding claims which additionally contains a pigment.
 - 13. A method of glazing a ceramic body, which method comprises coating a ceramic body with a glaze composition as claimed in any one of the preceding claims and firing the coated ceramic body at a glazing temperature of 950 to 1250°C.
- 14. A method as claimed in claim 13 wherein the ceramic body is ceramic table or whiteware including bone or fine china, vitreous and earthenware.
 - 15. A method as claimed in claim 13 or claim 14 wherein the ceramic body is coated by dipping or spraying.

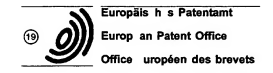
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- (54) Glaze compositions.
- GT A glaze composition comprises 45 to 75% of SiO₂, 0.1 to below 5% of Bi₂O₃, 0.1 to 20% of Al₂O₃, 2 to 20% of B₂O₃, at least one of CaO, MgO, SrO, BaO or ZnO in an amount of 2 to 22%, at least one of Li₂O, Na₂O, K₂O in an amount of 1 to 10%, at least one of La₂O₃, MoO₃ or WO₃ in an amount of 0.1 to 10% and certain other optional ingredients, all percentages being percentages by weight based on the total weight of the glaze composition, the glaze composition having a firing temperature of 950 to 1250°C, and having a coefficient of thermal expansion of 50 to 100 X 10⁻⁷/°C.



EUROPEAN SEARCH REPORT

Application Number

EP 92 30 3393

Category	Citation of document with of relevant p	indication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL5)
X	EP-A-0 267 154 (CI * page 1, line 32 * page 2, line 44	•	1-4,6, 8-15	C03C3/093 C03C3/095 C03C8/04 C03C8/14
٨	PATENT ABSTRACTS 0 vol. 13, no. 367 (& JP-A-01 122 937 1989 * abstract *	F JAPAN C-626)15 August 1989 (NAKASHIMA KK) 16 May	1	C04B41/86
A	DATABASE WPIL Week 9001, Derwent Publication AN 90-003955 & JP-A-01 286 937 November 1989 * abstract *	ns Ltd., London, GB; (NAKASHIMA KK) 17	1	
A	CHEMICAL ABSTRACTS 8 April 1991, Colum abstract no. 12770 page 322; * abstract * & JP-A-02 208 236 (17 August 1990	nbus, Ohio, US; 7p,	1	TECHNICAL FIELDS SEARCHED (Int. CL5)
A	FR-A-2 495 190 (EI: GMBH &CO.) * claims *	SENWERKE FRIED. DUKER	1-15	
D,P, A	EP-A-0 452 065 (COC	DKSON GROUP)	1-15	
	The present search report has	•		
В	Place of search ERLIN	Date of completion of the search 30 MARCH 1993		KUEHNE H.C.
X : part Y : part docu A : tech	ATEGORY OF CITED DOCUME cularly relevant if taken alone cularly relevant if combined with an ment of the same category notogical background written discourse	E : earlier patent d	locument, but publi date I in the application	ished on, or